AMENDMENTS TO THE CLAIMS

- 1. (Currently amended) An intranasal, transdermal, or intradermal dose form pharmaceutical composition comprising 0.5 ng to 20 μg desmopressin and a pharmaceutically acceptable carrier in a dosage form adapted for intranasal, transdermal, or intradermal administration sufficient to establish in which when administered to a patient in accordance with packaged instructions establishes a steady plasma/serum desmopressin concentration in the range of from about 0.1 picograms desmopressin per ml plasma/serum to about a maximum of 10.0 picograms desmopressin per ml plasma/serum and to-decrease decreases urine production.
- 2. (Cancelled)
- (Currently amended) The pharmaceutical composition dose form of claim 1 comprising from about 0.05 ug to about 10 ug desmopressin.
- (Currently amended) The pharmaceutical composition dose form of claim 1 comprising from about 0.1 µg to about 2 µg desmopressin.
- 5. (Cancelled)
- (Currently amended) The pharmaceutical composition dose form of claim 1 in a dosage form adapted for transdermal delivery for application to the skin comprising a patch, gel, cream, ointment, or iontophore.
- 7. (Currently amended) The pharmaceutical composition dose form of claim 1 adapted for transdermal administration for application to the skin comprising a an intradermal patch.
- 8. (Cancelled)
- (Currently amended) The pharmaceutical composition dose form of claim 1 in a dosage form sufficient to establish which establishes in a patient a steady plasma/serum desmopressin

concentration of from about 0.5 picograms desmopressin per ml plasma/serum to about 5.0 picograms desmopressin per ml plasma/serum.

10-26 (Cancelled)

- 27. (Currently amended) An pharmaceutical-intranasal dose form comprising desmopressin and a pharmaceutically acceptable carrier adapted for intranasal administration which when administered intranasally to a patient in accordance with packaged instructions establishes a steady plasma/serum desmopressin concentration in the range of from about 0.1 picograms desmopressin per ml plasma/serum to about a maximum of 10.0 picograms desmopressin per ml plasma/serum for a time between four and six hours and decreases urine production.
- 28. (Currently amended) The eomposition dose form of claim 27 which establishes in a patient a steady plasma/serum desmopressin concentration of from about 0.5 picograms desmopressin per ml plasma/serum to about 5.0 picograms desmopressin per ml plasma/serum.
- 29. (Currently amended) An intradermal or transdermal pharmaceutical dosage dose form comprising desmopressin and a pharmaceutically acceptable carrier for intranasal or transdermal administration which when administered intradermally or transdermally to a patient establishes a steady plasma/serum desmopressin concentration in the range of from about 0.1 picograms desmopressin per ml plasma/serum to about a maximum of 10.0 picograms desmopressin per ml plasma/serum for a time between four and six hours and decreases urine production.
- 30. (Currently Amended) The dosage dose form of claim 29 which establishes a steady plasma/serum desmopressin concentration of from about 0.5 picograms desmopressin per ml plasma/serum to about 5.0 picograms desmopressin per ml plasma/serum.
- 31. (Currently Amended) The dosage dose form of claim 29 comprising between $0.05~\mu g$ and $10~\mu g$ desmopressin.

- 32. (Currently Amended) The dosage dose form of claim 29 adapted for comprising an intradermal administration comprising a patch.
- 33. (Currently Amended) The dosage dose form of claim 29 adapted for transdermal delivery and comprising a patch, gel, cream, ointment, or iontophore.